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CYCLIC PEROXYACETAL COMPOSIDE

Technical Field

This invention relates to novel cyclic peroxyacetal compounds and in particular relates to compounds related to qinghaosu (Artemisinin) which is a naturally occurring biologically active compound.

Background of the Invention

The present invention provides novel compounds which are structurally similar to the naturally occurring biologically active compound qinghaosu (Artemisinin) which has the following formula:

Qinghaosu is a potent anti-malarial which has been successfully used to treat patients suffering of malaria re-emergence of strains The malaria. resistant to conventional (chloroquine) therapy is posing a world-wide problem and indeed, there is no universally acceptable cure at the present time. Qinghaosu occurs to the extent of about 0.1% (dry weight) in an annual shrub, qinghao or Artemisia annua, which grows in most provinces of China. Unfortunately, the world demand for qinghaosu considerable and there is supply, far exceeds the pressure to develop bioactive analogues and derivatives or to develop alternative sources for the compound.

Some of the novel compounds have activities superior to that of qinghaosu and furthermore these compounds will

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$$H_{3}C$$

$$(C)_{1}$$

$$(V)_{p}$$

$$(II)$$

wherein V is as defined for X; r and l are as hereinbefore defined; provided that the $(C)_{l}$ - $(V)_{r}$ group is not $CH(CH_{2})CH_{2}OH$ and $CH(CH_{3})C(=O)H$.

The present invention also provides pharmaceutical compositions comprising a compound of formula (I), a pharmaceutically acceptable salt thereof or stereoisomeric forms thereof in a pharmaceutically acceptable carrier and/or diluent.

Pharmaceutical compositions containing a compound of 10 formula (I) as the active ingredient in admixture with a pharmaceutically acceptable carrier or diluent can be pharmaceutical conventional according to prepared The carrier may be of any form formulating techniques. of preparation desired the form depending on 15 administration, eg intravenous, oral or parenteral.

In yet another aspect, the present invention provides a method of treatment or prophylaxis of parasitic or viral diseases in a mammal comprising administering to the mammal a compound of formula (I), a pharmaceutically acceptable salt thereof or a stereoisomeric form thereof.

Specific embodiments of the present invention are illustrated by the following preparative examples. It will be understood, however, that the invention is not confined to the specific limitations set forth in the individual examples.

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general, e residue after evaporation of the filtrate, without purification, was immediately submitted to Swern oxidation conditions. Thus, DMSO (0.37 ml; 5.2 mmol) in dichloromethane (2 ml) was added dropwise to a solution of oxalyl chloride (0.23 ml; 2.6 mmol) in dichloromethane (8 ml) at between -50° and -60° . After 5 min. the crude alcohol (288 mg) in dichloromethane (4 ml) was added dropwise at the same temperature and after 15 min. triethylamine (1.09 ml; 7.8 mmol) was added. was continued at -60° for 5 min. and then the whole was allowed to warm to room temperature over 20 min. workup afforded the aldehyde (2) which was submitted to flash chromatography (ether/light petroleum, give a colourless oil [228 mg; 79% from qinghao acid]. The aldehyde was found to be a 6.2:1 mixture of diastereomers and as such was used to prepare the other qinghao acid derivatives. ¹H NMR spectrum (200 MHz, CDCl₃) δ 0.877 (3H, d, J_{4-Me,4} = 6.3 Hz, 4-CH₃), 1.067 (3H, d, $J_{3',2'} = 7.0 \text{ Hz}$, H3'), 1.638 (3H, m, 7-CH₃), 5.132 (1H, m, H8), 9.588 (1H, d, $J_{1'.2'} = 4.2 \text{ Hz}$, H1').

Example 2

b) Ethyl alcohol (3)

The aldehyde (50.6 mg; 0.23 mmol) was dissolved in diethyl ether (3 ml) and treated with ethyl magnesium bromide in ether with cooling in an ice bath. The whole was then stirred at room temperature for 30 min. before being quenched with aqueous ammonium chloride solution (ice bath). Acidic workup gave the crude alcohol which was purified by flash chromatography (ether/light petroleum, 20:80) as a colourless viscous oil (48.8 mg;

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85%) and as a 85:15 mixture of epimers. v_{max} 3611 m, 3466 br m (OH), 3011 m, 2964 s, 2924 vs, 2872 s, 1456 m, 1380 m, 1236 w, 982 m, 951 m, cm⁻¹. ¹H NMR spectrum (200 MHz, CDCl₃) d 0.846 (3H, d, $J_{1',2'} = 6.5$ Hz, H1'), 0.868 (3H, d, $J_{4-Me,4} = 6.3$ Hz, $4-CH_3$), 0.955 (3H, t, $J_{5',4'} = 7.4$ Hz, H5'), 1.632 (3H, m, 7-CH₃), 2.479 (1H, br s, $W_{h/2} = 11.3$ Hz, OH), 3.736 (1H, m, H3'), 5.188 (1H, m, H8). Mass spectrum: m/z 250 (M, 2%), 232 (M-H₂O, 22), 189 (37), 162 (100), 81 (41).

10 Example 3

c) Phenyl alcohol (4)

The aldehyde (48 mg; 0.22 mmol) in ether (4 ml) was treated with phenyl magnesium bromide in ether as described above. Flash chromatography (ether/light petroleum, 20:80) of the crude product gave the phenyl alcohol (4) as a colourless viscous oil (55.8 mg; 86%). This was found to be a 76:24 mixture of epimers with respect to the hydroxyl group. Partial $^{l}{\rm H}$ NMR spectrum (200 MHz, CDCl3, *denotes minor epimer) δ *2.340 (1H, br s, Wh/2 = 11.3 Hz, OH), 2.512 (1H, br s, Wh/2=11.3 Hz, OH), *5.043 (1H, d, $J_{1^{\circ},2^{\circ}}=4.3$ Hz, H1'), 5.107 (1H, m, H1'), 5.171 (1H, m, H8), *5.277 (1H, m, H8).

Example 4

d) Allyl alcohol (5)

The alderyde (95.1 mg; 0.43 mmol) in ether (5 ml) was treated with allylmagnesium bromide in ether as described above. Flash chromatography (ether/light petroleum, 20:80) of the crude product gave the allyl alcohol (5) as a colourless viscous oil (99.7 mg; 88%) and as an approximately 55:45 mixture of epimers. ¹H NMR spectrum (200 MHz, CDCl₃, *denotes minor epimer) δ 0.865 (3H, d, $J_{4-Me,4} = J_{1',2'} = 6.2$ Hz, $4-CH_3$, H1'), *0.881 (3H, d, $J_{4-Me,4} = 6.5$ Hz, $4-CH_3$), *0.897 (3H, d, $J_{1',2'} = 6.7$ Hz, H1'), 1.62 (3H, m, 7-CH₃, *7-CH₃), 2.48 (1H, br s, $W_{h/2} = 12$ Hz, OH, *OH), 3.83-3.95 (1H, m, H3', *H3'), 5.07-5.29 (3H, m, 2 x H6', 2 x *H6', H8, *H8), 5.73-5.97 (1H, m, H5', *H5').

Example 5

e) Hydroxy acetals (6a) and (6b)

The aldehyde (210.7 mg; 0.96 mmol) in ether (8 ml) was 15 treated with the Grignard reagent (1.5 equiv.), derived from 5-bromopentanaldehyde diethyl acetal and magnesium in THF, with cooling in an ice bath. The whole was stirred at room temperature for 30 min. before being quenched with aqueous ammonium chloride. The product was 20. extracted into ether, and the ether extracts were washed with brine and dried (Na2SO4). Evaporation of the solvents under reduced pressure left a pale liquid which upon purification by flash chromatography (ether/light petroleum, 35:65) gave the hydroxy diethyl acetal as a 25 colourless viscous oil (309 mg; 85%) and as a ca. 88:12 mixture of the (2'R,3'R) - and (2'R,3'S) -diastereomers, (6a) and (6b) respectively. These were separated by h.p.l.c. (ethyl acetate/light petroleum, 13:87, Whatman

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Partisil 10, preparative) to give the $(2^{\circ}R, 3^{\circ}S)$ -isomer (6a) as the first to be eluted:- $[\alpha]_D^{20}$ +2.1° (c, 0.43, CHCl₃). v_{max} (CHCl₃) 3536 w, 3600-3100 br s (OH), 2977 s, 2931 s, 2870 s, 1454 m, 1377 m, 1236 w, 1126 s, 1057 s, 993 m cm⁻¹. ¹H NMR spectrum (200 MHz, CDCl₃) δ 0.845 (3H, d, $J_{1',2'} = 6.4$ Hz, H1'), 0.866 (3H, d, $J_{4-Me,4} = 6.3$ Hz, 4-CH₃), 1.205 (6H, t, J = 7.1 Hz, CH₃CH₂O), 1.631 (3H, m, 7-CH₃), 2.471 (1H, br s, $W_{h/2} = 11.5$ Hz, OH), 3.41-3.72 (4H, m, CH₃CH₂O), 3.82 (1H, m, H3'), 4.491 (1H, t, $J_{8',7'} = 5.6$ Hz, H8'), 5.180 (1H, m, H8). Mass spectrum: m/z 379 (M-1, 0.5%), 335 (10), 317 (11), 289 (16), 189 (47), 162 (100), 143 (43), 103 (66), 85 (52), 81 (61), 69 (53), 55 (66), 47 (57), 31 (62).

The next to be eluted was the $(2^{\circ}R, 3^{\circ}S)$ -isomer (6b):- $[\alpha]_D^{20}$ -7.3° (c, 0.30, CHCl₃). v_{max} (CHCl₃) 3535 w, 3600 - 3100 br s (OH), 2977 s, 2934 s, 2873 s, 1455 m, 1377 m, 1234 w, 1126 s, 1057 s, 1002 m cm⁻¹. 1H NMR spectrum (200 MHz, CDCl₃) δ 0.862 (6H, d, $J_{1',2'}$ = $J_{4-Me,4}$ = 6.6 Hz, H1', 4-CH₃), 1.207 (6H, t, J = 7.1 Hz, CH₃CH₂O), 1.625 (3H, m, 7-CH₃), 2.461 (1H, br s, $W_{h/2}$ = 11.2 Hz, OH), 3.42-3.73 (4H, m, CH₃CH₂O), 3.82 (1H, m, H3'), 4.492 (1H, t, $J_{8',7'}$ = 5.6 Hz, H8'), 5.244 (1H, m, H8). Mass spectrum: m/z 379 (M-1, <0.1%), 335 (1), 317 (2), 189 (17), 162 (100), 143 (15), 103 (52), 98 (27), 95 (19), 85 (23), 81 (37), 75 (21), 69 (21), 55 (32), 43 (25), 31 (37).

Example 6

f) Hydroxy acid (7a) and (7b)

The (2'R,3'R)-hydroxy acetal (6a) (40.4 mg; 0.11 mmol) was dissolved in THF (2 ml) and treated with aqueous HCl

1) with rapid stirring at room temperature. After 30 min. the whole was extracted with ether and the combined extracts were washed with brine and dried Evaporation of the solvents left the crude aldehyde which was then dissolved in ethanol (1 ml). This was added to a stirred solution of silver nitrate (54 mg; 0.32 mmol) in water (0.12 ml). The resulting mixture was cooled in an ice bath, treated with aqueous KOH solution (34%, 0.105 ml), and then stirred at room temperature for 2 h. The black silver precipitate was 10 filtered off, washing with water, and the filtrate was acidified with 3 M HCl with cooling in ice. The acidic was extracted with ether and the combined extracts were dried $(MgSO_4)$ and evaporated to dryness. Purification of the residue by chromatography on silicic 15 acid (ether/light petroleum, 1:1) afforded the (2'R,3'R)hydroxy acid (7a) as a colourless viscous oil (31.2 mg; 91% overall). ¹H NMR spectrum (200 MHz, CDCl₃) δ 0.864 (3H, d, $J_{4-Me,4} = 6.3$ Hz, $4-CH_3$), 0.881 (3H, d, $J_{1',2'} = 6.3$ Hz, H1'), 1.648 (3H, m, 7-CH₃), 2.390 (2H, t, $J_{7'.6'} = 7.3$ 20 Hz, H7'), 2.480 (1H, br s, $W_{h/2} = 12.8$ Hz, OH), 3.86 (1H, m, H3'), 5.185 (1H, m, H8), 5.969 (1H, br s, $W_{h/2}=88.5$ Hz, Similarly, the (2'R,3'S)-hydroxy acetal (6b) (30.4 mg; 80 μ mol) was converted into the (2'R,3'S)hydroxy acid (7b) (24 mg; 93%). ¹H NMR spectrum (200 MHz, 25 CDCl₃) δ 0.866 (3H, d, J_{1',2'} = J_{4-Me,4} = 6.8 Hz, H1', 4-CH₃), 1.630 (3H, m, 7-CH₃), 2.382 (2H, t, $J_{7',6'} = 7.3$ Hz, H7'), 2.461 (1H, br s, $W_{h/2} = 11.9 \text{ Hz}$, OH), 3.85 (1H, m, H3'), 4.435 (1H, br s, $W_{h/2}$ = 113 Hz, COOH), 5.238 (1H, m, H8).

30 Example 7

q) Dimethyl alcohol (8)

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Methylmagnesium iodide was prepared by treating magnesium turnings (80 mg; 3.3 mmol) in dry ether (4 ml) with methyl iodide (0.17 ml; 2.8 mmol) under gentle reflux. A solution of the methyl ester of dihydroginghao acid (138.3 mg; 0.55 mmol) in ether (3 ml) was then added dropwise at room temperature. The reaction mixture was heated under gentle reflux for a further 2 h before being cooled in ice. Aqueous ammonium chloride solution was added and the whole was extracted with ether. combined ether extracts were washed with brine, dried (Na₂SO₄) and evaporated to dryness to give the crude This was purified by flash chromatography alcohol. 15:85) to give the dimethyl (ether/light petroleum, alcohol as a crystalline solid (113 mg; 82%), m.p. 52-54°. ^{1}H NMR spectrum (200 MHz, CDCl3) δ 0.855 (3H, d, J4 $Me,4 = 6.4 \text{ Hz}, 4-CH_3), 0.926 (3H, d, <math>J_{1',2'} = 6.9 \text{ Hz}, \text{ H1'}),$ 1.190 (3H, s, H4' or 3-CH₃), 1.249 (3H, s, H4' or 3-CH₃), 0.87 - 2.0 (13H, m), 1.628 (3H, br m, $W_{h/2}$ = 5.5 Hz, 7- $\mathrm{CH_3}$), 2.526 (1H, br s, $\mathrm{W_{h/2}}$ = 11.6 Hz, OH), 5.280 (1H, br m, $W_{h/2} = 6.8 \text{ Hz}$, H8). Mass spectrum: m/z 232 (M - H₂O, 9%), 217 (7), 189 (25), 163 (36), 162 (100), 147 (17), 121 (13), 107 (17), 95 (14), 81 (27), 59 (40), 41 (20).

PREPARATION OF FINAL COMPOUNDS
Preparation of Deoxoqinghaosu Derivatives

25 Example 8

a) Deoxoqinghaosu (9)

Dihydroqinghao alcohol (1) (43.9 mg; 0.20 mmol) was mixed with acetonitrile (2.5 ml) and irradiated under oxygen at -30° for 3 h in the presence of Rose Bengal sensitiser. The resulting hydroperoxide solution was then diluted

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with dichroromethane (5 ml) and treated with Cu(OTf)2 (0.020 mmol, 0.1 M in acetonitrile) at -20° . reaction mixture was stirred at -150 for a further 40 min. and was then allowed to warm to room temperature over 20 min. before being cooled to -10°. added and then the whole was extracted with ether. combined extracts were washed with brine, dried (MgSO₄) The residue was purified by and evaporated to dryness. flash chromatography (ether/light petroleum, 40:60) to give deoxoqinghaosu (9) as a crystalline solid (19 mg; 36%). ¹H NMR spectrum (200 MHz, CDCl₃) δ 0.780 (3H, d, $J_{9-Me.9}=7.2 \text{ Hz}$, 9-CH₃), 0.963 (3H, d, $J_{6-Me.6}=5.9 \text{ Hz}$, 6-CH₃), 1.43 (3H, s, 3-CH₃), 3.449 (1H, dd, $J_{gem}=11.7$, $J_{10,9}=11.7$ Hz, H10), 3.731 (1H, br dd, $J_{gem}=11.7$, $J_{10.9}=4$ Hz, H10), 5.200 (1H, s, H12).

Example 9

b) Ethyl Deoxoginghaosu (10)

The ethyl alcohol (3) (77.9 mg; 0.31 mmol) was dissolved in dichloromethane (1 ml) and acetonitrile (2 ml) and irradiated under oxygen at -30° for 4 h in the presence of Rose Bengal sensitiser. The resulting solution of hydroperoxides was then diluted with dichloromethane (7 ml) and treated with Cu(OTf)₂ (0.031 mmol, 0.1 M in acetonitrile) at -20°. The mixture was kept at -20° for a further 40 min. and then allowed to warm to room temperature over 20 min. before being cooled to -10°. Water was added and then the whole was extracted into ether. The combined extracts were washed with brine, dried (MgSO₄) and evaporated to dryness. The residue was fractionated by flash chromatography (ether/light

petroleum, 10:90) to give ethyl deoxoginghaosu (10) as a waxy solid (31.1 mg; 34%) and essentially only as the Gepimer with respect to the ethyl group, $^{1}\mathrm{H}$ NMR spectrum (400 MHz, CDCl₃) δ 0.860 (3H, d, J_{Me,9}=7.6 Hz, 9-CH₃), 0.958 (3H, d, $J_{Me.6}=6.0$ Hz, 6-CH₃), 1.034 (3H, t, $J_{2'.1}=7.3$ Hz, H2), 1.426 (3H, s, 3-CH₃), 2.687 (1H, ddq, $J_{9.Me}=7.5$, $J_{9,8a}=7.5$, $J_{9,10}=6.1$ Hz, H9), 4.036 (1H, ddd, $J_{10,1}=10.6$, $J_{10,9}=6.0$, $J_{10,1}=2.8$ Hz, H10), 5.28 (1H, s, H12). ¹³C NMR spectrum (50 MHz, CDCl₃) δ 12.105 (C2'), 13.01 (9-CH₃), 20.20 (6-CH₃), 22.27 (C8), 24.87 (C1' or C5), 24.87 (C1' 10 or C5), 26.13 (3-CH₃), 30.25 (C9), 34.48 (C7), 36.57 (C4), 37.42 (C6), 44.45 (C8a), 52.03 (C5a), 77.64 (C10), 81.14 (C12a), 88.88 (C12), 103.14 (C3). Mass spectrum: m/z 296 (M⁺, 0.7%), 278 (M-H₂O, 1), 264 (M-O₂, 24), 206 (100), 193 (33), 182 (45), 165 (35), 124 (72), 95 (35), 15 81 (34), 69 (58), 55 (92), 43 (90), 41 (63), 29 (38).

Example 10

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c) Phenyl Deoxoqinghaosu (11)



The phenyl alcohol (4) (55.4 mg; 0.19 mmol) was dissolved in dichloromethane (0.5 ml) and acetonitrile (1.5 ml) and photooxygenated at -30° for 3 h. Dilution of the reaction mixture with dichloromethane and treatment with ${\rm Cu}({\rm OTf})_2$ as described above followed by purification of the crude product by flash chromatography (ether/light petroleum, 7:93) afforded phenyl deoxoqinghaosu (11) as a viscous oil (17.4 mg; 27%) and exclusively as the ßepimer with respect to the phenyl group, $^1{\rm H}$ NMR spectrum (200 MHz, CDCl₃) δ 0.508 (3H, d, $J_{\rm Me,9}$ =7.7 Hz, 9-CH₃), 0.987 (3H, d, $J_{\rm Me,6}$ =5.6 Hz, 6-CH₃), 1.383 (3H, s, 3-

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CH₃),2.74. (1H, ddq, $J_{9,Me}=7.1$, $J_{9,8a}=7.1$, $J_{9,10}=6.7$ Hz, H9), 5.577 (1H, s, H12), 5.723 (1H, d, $J_{10,9}=6.7$ Hz, H10, 7.15-7.34 (5H, m, $C_{6}H_{5}$). ¹³C NMR spectrum (50 MHz, CDCl₃) d 13.75 (9-CH₃), 19.98 (6-CH₃), 24.84 (C5 or C8), 25.01 (C5 or C8), 25.81 (3-CH₃), 32.21 (C9), 34.29 (C7), 36.77 (C4), 37.60 (C6), 43.59 (C8a), 51.59 (C5a), 73.14 (C10), 81.26 (C12a), 90.95 (C12), 102.38 (C3), 126.23, 126.38, 127.80 ($C_{6}H_{5}$ C_{meta}, C_{para}, C_{ortho}), 141.15 ($C_{6}H_{5}$ C_{ipso}). Mass spectrum: m/z 326 (M⁺-H₂O, 0.4%), 312 (M-O₂, 13), 298 (12), 254 (16), 240 (23), 182 (100), 124 (68), 118 (52), 105 (31), 91 (32), 55 (23), 43 (46), 28 (61).

Example 11

d) Allyl Deoxoqinghaosu (12)

The allyl alcohol (5) (46 mg; 0.18 mmol) was submitted to a). Allyl described in the same conditions as 15 then obtained by flash (12)was deoxoginghaosu chromatography (ether/light petroleum, 10:90) crystalline solid (18.3 mg; 36%) and as a 1:3 mixture of α - and ß-epimers with respect to the allyl group, ^1H NMR spectrum (200 MHz, CDCl₃) δ 0.782 [3H, d, J_{Me.9}=7.2 Hz, 9-20 $CH_3(\alpha)$], 0.888 [3H, d, $J_{Me,9}=7.6$ Hz, 9- $CH_3(\beta)$], 0.951 [3H, d, $J_{\text{Me.6}}=6.0$ Hz, $6-\text{CH}_3(\alpha)$], 0.965 [3H, d, $J_{\text{Me.6}}=5.8$ Hz, 6- $CH_3(B)$], 1.417 (3H, s, 3-CH₃), 2.69 [1H,ddq, $J_{9.8a}=7.9$, $J_{9.Me}=7.3$, $J_{9.10}=6.3$ Hz, H9(B)}, 3.491 [1H, ddd, $J_{10.1}=10.2$, [1H, Hz, H10(α)], 4.304 $J_{10.9}=6.0$, $J_{10.1}=3.7$ 25 $J_{10,1}=10.0$, $J_{10,9}=6.1$, $J_{10,1}=4.3$ Hz, H10(G)], 5.02-5.18 (2H, m, H3'), 5.249 [1H, s, H12(α)], 5.332 [1H, s, H12(β)], 5.73-6.17 (1H, m, H2'). 13 C NMR spectrum (50 MHz, CDCl₃) δ

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be suitable for preparing conjugate drugs for the treatment of malaria and other parasitic and viral diseases.

The novel compounds can also be used as building blocks because of their reactive side chain and other active drugs can be linked via these side chains to form conjugate drugs.

Disclosure of the Invention

In one aspect, the present invention provides compounds of general formula (I), pharmaceutically acceptable salts thereof or stereoisomeric forms thereof

$$H_3C \xrightarrow{O-O} \begin{pmatrix} CH_3 \\ (\times)_n \end{pmatrix}$$

wherein

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1 = 1, 2 or 3

n is an integer from 1-6

where X is independently selected from

H, =0,=CH₂, aryl, COR, OR, COOR or

 $X = -(CR_1R_2)_r-R_3$ where r is an integer from 1-10 and where r > 1, optionally at least 1 carbon atom can be replaced by 0, S or N;

 R_1 , R_2 and R_3 are independently selected from

H; alkyl, alkenyl, alkynyl, aryl, each optionally substituted by one or more substituents selected from alkyl, alkenyl, alkynyl, aryl, halogen, OR, CF3, NO2, COOR, NRR', SR, COR, CONRR', SO3R, SO2NRR', SR, SOR, and

12.99 [9-CH₃(\mathbb{B})], 13.60 [9-CH₃(α)], 20.20 [6-CH₃(\mathbb{B})], 20.33 [6-CH₃(α)], 21.51 (CH₂), 24.72 (CH₂), 24.90 (CH₂), $(3-CH_3)$, 30.21 [C9(B)], 31.03 [C9(α)], [C7(B)], 34.49 [C7(α)], 36.35 [C4(α)], 36.61 [C4(B)], [C6(B)], [C6(α)], 37.49 (CH_2) , 37.40 37.16 5 [C8a(\mathfrak{L})], 46.07 [C8a(α)], 52.00 [C5a(α)], 52.36 [C5a(\mathfrak{L})], 73.87 [C10(α)], 74.73 [C10(β)], 80.70 [C12a(α)], 81.08 103.14 [C12(B)], 92.02 $[C12(\alpha)]$, 89.10 [C12a(B)], 116.42 [C3'(B)], 116.08 [C3(α)], 103.97 [C3(B)], [C3'(α)], 134.92 [C2'(α)], 136.48 [C2'(β)]. 10

Example 12

e) Carboxybutyl Deoxoqinghaosu (13a), (13b)

The hydroxy acid (7a) (30.2mg; 93.7 μ mol) was submitted to the same conditions as described in a). The resulting crude product was then fractionated by chromatography on 15 silicic acid (ether/light petroleum, 50:50) to give the G-epimer of carboxybutyl deoxoqinghaosu (13a) colourless viscous oil (11.9 mg; 34%). ¹H NMR spectrum (200 MHz, CDCl₃) δ 0.857 (3H, d, J_{9-Me,9}=7.6 Hz, 9-CH₃), 0.961 (3H, d, $J_{6-Me.6}=5.8$ Hz, 6-CH₃), 1.419 (3H, s, 3-CH₃), 20 4.150 (1H, ddd, $J_{10,1}$ =10.1, $J_{10,9}=6.2$, $J_{10,1}$ =2.2 Hz, H10), 5.298 (1H, s, H12). ^{13}C NMR spectrum (50 MHz, CDCl₃) δ 13.01 (9-CH₃), 20.19 (6-CH₃), 24.63, 24.63, 24.86 (C1', C5, C8), 26.10 (3-CH₃), 27.03, 29.01 (C3', C2'), 30.26 (C9), 33.88 (C4'), 34.45 (C7), 36.57 (C4), 37.44 (C6), 25 44.39 (C8a), 52.36 (C5a), 75.37 (C10), 81.13 (C12a), 88.99 (C12), 103.22 (C3), 178.98 (C5). Similarly, the hydroxy acid (7b) (18.3 mg; 57 μ mol) was converted into the α -epimer of carboxybutyl deoxoqinghaosu (13b) (8.4

mg; 40%). ¹H NMR spectrum (200 MHz, Cbcl₃) d 0.777 (3H, d, $J_{9-Me,9}=7.4$ Hz, 9-CH₃), 0.953 (3H, d, $J_{6-Me,6}=6.2$ Hz, 6-CH₃), 1.409 (3H, s, 3-CH₃), 3.42 (1H, m, H10), 5.219 (1H, s, H12).

5 Example 13

f) Dimethyl Deoxoqinghaosu (14)

The dimethyl alcohol (8) (53 mg; 0.21 mmol) was submitted to the same conditions as described in a). Dimethyl by flash isolated was then (14)deoxoginghaosu chromatography on silica (ether/light petroleum, 20:80) 10 as a viscous oil which slowly crystallised (22 mg; 35%). $^{\rm I}$ H NMR spectrum (200 MHz, CDCl₃) δ 0.887 (3H, d, J_{9-Me.9} = 7.5 Hz, 9-CH₃), 0.959 (3H, d, $J_{6-Me.6} = 5.8$ Hz, 6-CH₃), 1.228 (3H, s, 10-CH₃), 1.325 (3H, s, 10-CH₃), 1.424 (3H, s, 3-CH₃), 1.36-1.93 (9H, m), 2.027 (1H, ddd, J = 14.4, J15 = 4.7, J = 3.1 Hz), 2.277 (1H, ddd, J = 13.9, J = 13.4, J= 4.0 Hz), 2.433 (1H, dq, $J_{9,9-Me}$ = 7.5, $J_{9,8a}$ = 4.3 Hz, H9), 5.313 (1H, s, H12). 13 C NMR spectrum (50 MHz, CDCl₃) δ 13.96 (9-CH₃), 20.41 (6-CH₃), 20.76 (10-CH₃), 23.81 (C8), 24.56 (C5), 26.32 (3-CH₃), 31.91 (C9), 34.76 (C7), 20 35.68 (10-CH₃), 36.58 (C4), 37.48 (C6), 46.28 (C8a), 52.80 (C5a), 76.3 - 77.6 (C10, obscured by CDCl₃), 80.97 (C12a), 88.94 (C12), 103.84 (C3). Mass spectrum: m/z 264 $(M-O_2, 7\%)$, 250 (28), 235 (17), 207 (17), 206 (17), 192 (73), 177 (100), 165 (32), 138 (48), 123 (24), 109 (24), 25 95 (28), 81 (23), 69 (37), 55 (52), 43 (95), 28 (21).

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Preparation of Dehydroqinghaosu Adducts

<u>Example 14</u>

a) With thiophenol

A solution of dehydroqinghaosu (17.6 mg; 62.7 μ mol) in chloroform (2 ml) was treated with thiophenol (6.5 μ l; 5 62.7 μ mol) and triethylamine (4 μ l) with cooling in an ice bath. The whole was then stirred at room temperature under nitrogen for 24 h. Aqueous workup afforded the phenylthio derivative (15) as a 1:2 mixture of $\mathfrak B$ - and α epimers in essentially quantitative yield and free of by-10 products (23.9 mg; 98%). ¹H NMR spectrum (600 MHz, CDCl₃) δ 0.992 [3H, d, $J_{\text{Me},6}=6.3$ Hz, $6-\text{CH}_3(\alpha)$], 1.007 [3H, d, $J_{\text{Me},6}=5.9$ Hz, 6-CH₃(B)], 1.419 [3H, s, 3-CH₃(B)], 1.451 [3H, s, 3-CH₃(α)], 2.272 [1H, ddd, J_{9,1}=9.4, J_{9,1}=3.4, $J_{9.8a}=1$ Hz, H9(α)], 2.834 [1H, dd, $J_{gem}=13.7$, $J_{1,9}=11.8$ Hz, 15 H1'(B)], 3.139 [1H, dd, $J_{gem}=13.8$, $J_{1',9}=11.7$ Hz, H1'(α)], 3.446 [1H, ddd, $J_{9,1}=11.8$, $J_{9,1}=5$, $J_{9,8a}=5$ Hz, H9(£)], 3.776 [1H, dd, $J_{gem}=13.8$, $J_{1',9}=4.4$ Hz, H1'(B)], 3.860 [1H, dd, $J_{gem}=13.8$, $J_{1,9}=3.4$ Hz, H1'(α)], 5.864 [1H, s, H12(β)], 5.918 [1H, s, H12(α)], 7.18-7.40 (5H, m, SPh). 20

Example 15

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b) With thioglycolic acid.

A solution of dehydroqinghaosu (14.4 mg; 51.4 μ mol) in chloroform (2 ml) was treated with thioglycolic acid (3.6 μ l; 51.4 μ mol) and triethylamine (7.2 μ l; 51.4 μ mol).

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The whole was stirred at room temperature under nitrogen for 3 d before being quenched with NaHCO3 solution (0.3 The resulting mixture was extracted with ether and then the aqueous phase was acidified (HCl, 0.1M) to pH 6. Extraction of the aqueous phase with ether followed by the usual workup afforded the adduct (16) as a 2:1 mixture of g- and α -epimers (15.4 mg; 81%). These were separated by chromatography on silicic acid (ether/light petroleum, 60:40) as very hygroscopic fine white solids. First was obtained the G-epimer. ^{1}H NMR spectrum (200 MHz, CDCl₃) δ 1.007 (3H, d, J_{6-Me,6} = 5.6 Hz, 6-CH₃), 1.447 (3H, s, 3-CH₃), 5.878 (1H, s, H12). Next was obtained the α -epimer. ¹H NMR spectrum (200 MHz, CDCl₃) δ 1.004 (3H, d, $J_{6-Me.6} = 5.6$ Hz, $6-CH_3$), 1.449 (3H, s, 3-CH₃), 5.956 (1H, s, H12).

Example 16

c) With methyl thioglycolate.

Dehydroqinghaosu (16.7 mg; 59.5 μ mol) was treated with methyl thioglycolate (5.3 59.5 and $\mu l;$ μ mol) triethylamine (4 μ l) in chloroform (2 ml) for 3 d as described above. The reaction mixture was evaporated to dryness and then fractionated by flash chromatography (ether/light petroleum, 50:50) to give the ß-epimer as the least polar fraction (13.8 mg; 60%) and as very fine needles, ^{1}H NMR spectrum (200 MHz, CDCl₃) δ 1.00 (3H, d, $J_{\text{Me.6}=5.9}$ Hz, 6-CH₃), 1.44 (3H, s, 3-CH₃), 2.65 (1H, dd, $J_{gem}=13.2$, $J_{1',9}=10.6$ Hz, H1'), 3.21 (1H, d, $J_{gem}=14.9$ Hz, H1"), 3.32 (1H, d, $J_{gem}=14.9$ Hz, H1"), 3.40 (1H, dd, $J_{gem}=13.1$, $J_{1'.9}=5.0$ Hz, H1'), 3.51 (1H, ddd, $J_{9.1}=10.8$, $J_{9,1}$ =5.0, $J_{9,8a}$ =5 Hz, H9), 3.757 (3H, s, OCH₃), 5.871 (1H,

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s, H12), and the α -epimer (6.4 mg; 28%) as a viscous oil, $I_{\rm H}$ NMR spectrum (200 MHz, CDCl₃) δ 1.00 (3H, d, $J_{\rm 6-Me,6}$ = 5.3 Hz, 6-CH₃), 1.45 (3H, s, 3-CH₃), 1.2 - 2.5 (11H, m), 2.39 (1H, ddd, $J_{\rm 9,1}$ = 11.2, $J_{\rm 9,1}$ = 4.0, $J_{\rm 9,8a}$ = 1.2 Hz, H9), 3.00 (1H, dd, $J_{\rm gem}$ = 13.3, $J_{\rm 1,9}$ = 11.3 Hz, H1'), 3.25 (1H, d, $J_{\rm gem}$ = 14.9 Hz, H1"), 3.32 (1H, d, $J_{\rm gem}$ = 14.9 Hz, H1"), 3.41 (1H, dd, $J_{\rm gem}$ = 13.2, $J_{\rm 1,9}$ = 3.9 Hz, H1'), 3.74 (3H, s, OCH₃), 5.95 (1H, s, H12).

Example 17

10 d) With mercaptopropionic acid

A solution of dehydroqinghaosu (24.9 mg; 89 μ mol) in chloroform (2 ml) was treated with mercaptopropionic acid (7.8 μ l; 89 μ mol) and triethylamine (12.4 μ l; 89 μ mol) with stirring at room temperature under nitrogen for 24 h and then for a further 24 h allowing the solvent to slowly evaporate away. The whole was diluted with ether and washed with dilute HCl solution. The ether solution was dried (MgSO₄) and evaporated to dryness to give the crude adduct which was submitted to chromatography on silicic acid (ether/light petroleum, 75:25). The first compound to be eluted was the ß-epimer which was obtained as a fine white solid (8.2 mg; 24%). ¹H NMR spectrum (600 MHz, CDCl₃) δ 1.008 (3H, d, $J_{6-Me,6} = 5.7$ Hz, 6-CH₃), 1.041 - 1.155 (2H, m), 1.40 - 1.51 (4H, m), 1.445 (3H, s, $3-CH_3$), 1.76 - 1.83 (2H, m), 2.141 (1H, ddd, J = 13.0, J = 4.6, J = 4.6 Hz), 2.42 - 2.47 (1H, m), 2.554 (1H, dd, $J_{gem} = 13.5$, $J_{1,9} = 11.9$ Hz, H11), 2.67 - 2.70 (2H, m, 2 X H2"), 2.79 - 2.81 (2H, m, 2 X H1"), 3.334 (1H, dd, $J_{gem} =$ 13.6, $J_{1'.9} = 4.6 \text{ Hz}$, H1'), 3.449 (1H, ddd, $J_{9.1'} = 11.6$,

 $J_{9,1'}=4.6$, $J_{9,8a}=4.8$ Hz, H9), 5.865 (1H, s, H12). Next was obtained the α -epimer as a viscous oil (24.5 mg; 71%). ¹H NMR spectrum (200 MHz, CDCl₃) δ 1.060 (3H, d, $J_{6-Me,6}=5.8$ Hz, 6-CH₃), 1.506 (3H, s, 3-CH₃), 1.15 - 2.61 (11H, m, 2 X H4, 2 X H5, H5a, H6, 2 X H7, 2 X H8, H8a), 2.355 (1H, ddd, $J_{9,1'}=11.5$, $J_{9,1'}=3.7$, $J_{9,8a}=1.0$ Hz, H9), 2.70 - 2.90 (4H, m, 2 X H1", 2 X H2"), 2.952 (1H, dd, $J_{1',1'}=13.3$, $J_{1',9}=11.6$ Hz, H1'), 3.401 (1H, dd, $J_{1',1'}=13.3$, $J_{1',9}=3.7$ Hz, H1'), 5.96 (1H, s, H12).

10 Example 18

e) With methyl mercaptopropionate

Dehydroqinghaosu (29.4 mg; 0.105 mmol) in chloroform (3 ml) was treated with methyl mercaptopropionate (11.6 μ l; 0.105 mmol) and triethylamine (4 μ l) with stirring as 15 described above. After 48 h the whole was evaporated to dryness and the residue was submitted chromatography (ether/light petroleum, 50:50) to give first the ß-epimer as a fine white solid (18.3 mg; 44%). ¹H NMR spectrum 200 MHz, CDCl₃) δ 1.01 (3H, J_{6-Me.6} = 5.8 20 Hz, $6-CH_3$), 1.45 (3H, s, $3-CH_3$), 1.0- 1.9 (6H, m), 2.0 -2.2 (3H, m), 2.4 - 2.5 (2H, m), 2.54 (1H, dd, $J_{gem} = 13.2$, $J_{1',9} = 11.4 \text{ Hz}, \text{ H1'}, 2.59 - 2.67 (2H, m, 2 X H2"), 2.76 -$ 2.84 (2H, m, 2 X H1"), 3.33 (1H, dd, $J_{gem} = 13.3$, $J_{1'.9} =$ 4.7 Hz, H1'), 3.44 (1H, ddd, $J_{9,1}' = 11.5$, $J_{9,1}' = 4.9$, $J_{9,8a}$ = 4.9 Hz, H9), 3.71 (3H, s, OCH₃), 5.67 (1H, s, H12). 13 C NMR spectrum (50 MHz, CDCl₃) δ 19.76 (6-CH₃), 22.97 (C7), 24.84 (C8), 25.12 (3-CH₃), 26.98 (CH₂), 28.86 (CH₂), 33.30 (CH₂), 34.41 (CH₂), 35.83 (CH₂), 37.51 (C6), 37.55 (C8a),

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41.17 (C5a), 49.97 (C9), 51.83 (OCH₃), 79.24 (C12a), 93.80 (C12), 105.49 (C3), 170.17 (COOCH₃), 172.08 (C10). Next was obtained the α-epimer as a viscous oil (9.7 mg; 23%). ¹H NMR spectrum (200 MHz, CDCl₃) δ 1.00 (3H, d, J₆. 5 Me,6 = 5.7 Hz, 6-CH₃), 1.1 - 1.6 (5H, m), 1.45 (3H, s, 3-CH₃), 1.65 -1.83 (2H, m), 1.69 -2.08 (2H, m), 2.14 (1H, br dd, J_{gem} = 13.7, J = 4.5 Hz), 2.29 (1H, ddd, J_{9,1} = 11.6, J_{9,1} = 3.7, J_{9,8a} = 1.1 Hz, H9), 2.3 - 2.47 (1H, m), 2.59 - 2.67 (2H, m, 2 X H2"), 2.76 - 2.84 (2H, m, 2 X H1"), 2.88 (1H, dd, J_{gem} = 13.2, J₁,9 = 3.7 Hz, H1'), 3.70 (3H, s, OCH₃), 5.95 (1H, s, H12). When the solvent was changed to dichloromethane the β-epimer was obtained in only 15% yield and the α-epimer in 77% yield.

15 Example 19

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f) With thioglycerol

Dehydroqinghaosu (14.6 mg; 52 μ mol) in chloroform (2 ml) was treated with thioglycerol (5.6 mg; 4.3 μ l; 52 μ mol) and triethylamine (4 μ l). The whole was stirred at room temperature overnight after which time complete reaction had taken place. The reaction mixture was evaporated to dryness to give the crude £- and α -epimeric adducts in a ratio of 2.2 : 1. These were separated by flash chromatography on silica (ethyl acetate/light petroleum, 80 : 20) to give first, the £-epimer as a mixture of diastereomers at C2" and as a fine white solid (12.2 mg; 60%). 1 H NMR spectrum (400 MHz, CDCl₃) (*denotes other diastereomer) δ 1.012 (3H, d, J₆-Me,6 = 5.7 Hz, 6-CH₃), 1.07 - 1.14 (2H, m), 2.00 - 2.18 (3H, m), 2.40 - 2.48

54 - 2.74 (3H, m), 3.320 (1H, add, J = 13.1, J)(1H, m), = 7.9, J = 4.8 Hz), *3.473 (1H, ddd, $J_{9.1}$ = 11.2, $J_{9.1}$ = 5.9, $J_{9.8a} = 3.6 \text{ Hz}$, H9), 3.482 (1H, ddd, $J_{9.1} = 10.9$, $J_{9.1}$ = 5.4, $J_{9.8a}$ = 5.4 Hz, H9), *3.572 (1H, dd, J = 11.2, J = 5.8 Hz), 3.580 (1H, dd, J = 11.2, J = 5.9 Hz), 3.758 (1H,5 br ddd, J = 11.3, J = 3.0, J = 3.0 Hz), 3.82 -3.88 (1H, m), 5.878 (1H, s, H12). 13 C NMR spectrum (50 MHz, CDCl₃) δ 19.79 (6-CH₃), 23.17 (C8), 24.85 (C5), 25.14 (3-CH₃), 28.92, 29.72 (CH_2) , 30.94 (C9), 33.35 (C7), 35.87 (C6), 37.53 (CH), 37.80, 38.19 (CH), 41.52, 41.73 (CH), 50.01 10 (C5a), 65.29 (C3"), 69.74, 70.37 (C2"), 79.29 (C12a), 93.89 (C12), 105.59 (C3), 170.25 (C10). The α -epimer was obtained as a viscous oil and as a mixture of diastereomers at C2" (5.1 mg; 25%). H NMR spectrum (400 MHz, CDCl₃) δ 1.009 (3H, d, $J_{6-Me,6} = 6.1$ Hz, 6-CH₃), 1.16 15 - 1.34 (2H, m), 1.39 - 1.52 (2H, m), 1.452 (3H, s, 3- CH_3), 1.70 - 1.82 (2H, m), 1.92 - 2.16 (3H, m), 2.31 -2.43 (2H, m), 2.59 - 2.75 (3H, m), 2.933 (1H, ddd, J =14.0, J = 11.1, J = 3.1 Hz), 3.329 (1H, ddd, J = 13.2, J= 13.2, J = 4.0 Hz), 3.55 - 3.60 (1H, m), 3.766 (1H, br 20 d, J = 11.4 Hz), 3.81 - 3.89 (1H, m), 5.954 (1H, s, H12).

Example 20

g) With L-cysteine

L-Cysteine (5.8 mg; 36.8 μ mol) was added as a solid to a stirred solution of dehydroqinghaosu (10.3 mg; 36.8 μ mol) in methanol (1.5 ml) under nitrogen. Complete conversion to the adduct occurred almost immediately. The reaction mixture was then evaporated to dryness whereupon the crude residue was triturated with ether and the resulting

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fine white powder was obtained by filtration. The adduct (17), so obtained, was found to be only sparingly soluble and insoluble in most other solvents. in methanol Therefore, an estimate of the epimer ratio could not be made. ^{1}H NMR spectrum (200 MHz, CD₃OD) δ 0.87 [3H, d, $J_{\text{Me},6}=6.4$ Hz, 6-CH₃(B)], 1.006 [3H, d, $J_{6-\text{Me}}=5.8$ Hz, 6- $CH_3(\alpha)$], 1.390 [3H, s, 3- $CH_3(\beta)$], 1.396 [3H, s, 3- $CH_3(\alpha)$], 2.821 [1H, dd, $J_{1',1'} = 12.9$, $J_{1',9} = 10.0$ Hz, H1'(B)], 2.951 [1H, dd, $J_{2",2"} = 14.6$, $J_{2",1"} = 8.5$ Hz, $H_{2"}(\alpha)$], 2.966 [1H, dd, $J_{2",2"} = 14.4$, $J_{2",1"} = 8.5$ Hz, H2"(£)], 3.167 [1H, dd, $J_{2",2"} = 14.6$, $J_{2",1"} = 4.0$ Hz, H2"(α)], 3.191 [1H, dd, $J_{2",2"} = 4.0$ 14.4, $J_{2^{"}.1^{"}} = 4.0 \text{ Hz}$, $H_{2^{"}}(\mathfrak{E})$], 3.490 [1H, ddd, $J_{9.1} = 10.0$, $J_{9,1} = 5.4$, $J_{9,8} = 5.4$ Hz, H9(E)], 3.702 [1H, dd, $J_{1},2 = 8.6$, $J_{1",2"} = 4.0 \text{ Hz}$, $H_{1"}(S)$], 3.749 [1H, dd, $J_{1",2"} = 8.4$, $J_{1",2"} = 8.4$ 3.7 Hz, $H1^{\alpha}(\alpha)$], 6.026 [1H, s, $H12(\beta)$], 6.127 [1H, s, $H12(\alpha)$

Example 21

h) With N-acetyl-L-cysteine

Triethylamine (7.9 μ l; 57 μ mol) was added to a stirred solution of dehydroqinghaosu (15.9 mg; 57 μ mol) and a suspension of N-acetyl-L-cysteine (9.3 mg; 57 μ mol) in chloroform (3 ml). Deprotonation of the cysteine caused it to go into solution. The reaction mixture was stirred under nitrogen for 24 h and then for a further 24 h allowing the solvent to evaporate away slowly. Water was added and then the whole was extracted once with ether. The aqueous layer was separated and then acidified with 1 M hydrochloric acid in the presence of ether and extracted three times with more ether. The combined

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extracts were dried (MgSO₄) and evaporated to dryness to give the cysteine adducts as a 1:1 mixture of α - and ß-epimers and as a viscous oil (18.9 mg; 75%). ¹H NMR spectrum (200 MHz, CDCl₃) δ 1.00 (3H, d, J_{6-Me,6} = 5.4 Hz, 6-CH₃), 1.44 (3H, s, 3-CH₃), 2.10 (3H, s, acetyl), 4.82 (1H, br m, W_{h/2} = 15.7 Hz, N-H), 5.875 [1H, s, H12(ß)], 5.961 [1H, s, H12(α)], 6.822 (1H, dd, J_{2",1"} = 7.9, J_{2",1"} = 7.9 Hz, H2"), 7.45 (1H, br s, W_{h/2} = 28.6 Hz, COOH).

Preparation of Ring-Contracted Analogues of Qinghaosu 10 Derivatives

Example 22

a) Oxidative degradation of dihydroqinghao aldehyde (2)

A mixture of dihydroqinghao aldehyde (2) (105 mg; 0.48 mmol), DABCO (26 mg), cupric acetate monohydrate (50 mg) and 2,2'-bipyridyl (50 mg) in DMF (7 ml) was stirred for 12 h at $70-75^{\circ}$ and then the solvent was removed under vacuum. The resulting residue was submitted to flash chromatography (ether/light petroleum, 1:5) to afford the ketone (23) as a colourless viscous oil (77 mg; 78%) $[\alpha]_D^{18}$ -44.4° (c, 2.03, CHCl₃). H NMR spectrum (400 MHz, CDCl₃) δ 0.94 (3H, d, J=6 Hz), 1.65 (3H, s), 2.20 (3H,s), 2.95 (1H, br s), 4.48 (1H, s).

Example 23

b) Reduction

so₂R, where R and R' are independently selected from H, optionally substituted alkyl, optionally substituted alkynyl, optionally substituted aryl or optionally substituted arylalkyl wherein the optional substituents are as defined above.

Preferred compounds of formula (I) have the following structural formulae:

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The ketone (23) (98 mg; 0.47 mmol) in methanol (3 ml) was treated with NaBH4, portionwise, at 00 until a t.l.c. monitor showed that the reaction was complete. Water was added and the whole was thoroughly extracted with ether. The combined ether extracts were washed with brine and dried (Na2SO4) and evaporated to dryness. The residue was submitted to flash chromatography (ether/light petroleum, 2:8) to give the alcohols, (24a) and (24b) as a colourless viscous oil (98 mg; 99%) and as a 1:4 mixture of epimers. These were separated by h.p.l.c. (ethyl acetate/light petroleum, 8:92, Whatman partisil 10 M9, semi-preparative) to give, firstly, epimer (24b) $[\alpha]_D^{18}$ -10.3° (c, 2.34, CHCl₃). ¹H NMR spectum (400 MHz, CDCl₃) δ 0.88 (3H, d, J=6 Hz), 1.24 (3H, d, J=6 Hz), 1.63 (3H, br s), 2.72 (1H, br s), 3.81 (1H, dq), 5.34 (1H, s). The next to be eluted was epimer (24a) $[\alpha]_D^{18}$ -8.5° (c, 1.68, CHCl₃). 1 H NMR spectrum (400 MHz, CDCl₃) δ 0.89 (3H, d, J=6 Hz), 1.26 (3H, d, J=6.4 Hz), 1.62 (3H, br s), 2.46 (1H, br s), 3.775 (1H, dq), 5.14 (1H, s).

20 Example 24

c) Oxygenation to deoxoqinghaosu analogues (25a) and (25b)

The major alcohol epimer (24b) (55.8 mg; 0.27 mmol) in acetonitrile (2 ml) and dichloromethane (1 ml) was irradiated under oxygen at -30° for 4 h in the presence of Rose Bengal sensitiser. The resulting hydroperoxide solution was then diluted with dichloromethane (7 ml) and treated with $\text{Cu}(\text{OTf})_2$ (0.027 mmol, 0.1 M in acetonitrile) as described for the preparation of the

deoxoging aosu derivatives. The crude product mixture submitted to flash work-up obtained after was chromatography (ether/light petroleum, 1:6) to give the five membered ring analogue (25b) as a colourless viscous oil (19.1 mg; 28%) $[\alpha]_D^{20}+120.2^{\circ}$ (c, 1.1, CHCl₃). ¹H NMR 5 spectrum (400 MHz, CDCl₃) δ 0.98 (3H, d, J=6 Hz), 1.45 (3H, s), 1.47 (3H, d, J=6.5 Hz), 3.93 (1H, dq), 5.59 (1H, dq)Similarly, the minor alcohol epimer (24a) (20 mg; 96 μ mol) was converted into the five membered analogue (25a) (7.8 mg; 32%). ¹H NMR spectrum (400 MHz, CDCl₃) δ 0.98 10 (3H, d, J=6 Hz), 1.24 (3H, d, J=6.4 Hz), 1.44 (3H, s), 4.92 (1H, dq), 5.67 (1H,s).

Activity data

The testing results for the qinghaosu derivatives are as follows:

Ethyl deoxoqinghaosu has been tested against two strains of *Plasmodium falciparum* concurrently with artemether and sodium artesunate, the latter two compounds being in clinical use at the present time. The results are shown in the following graphs.

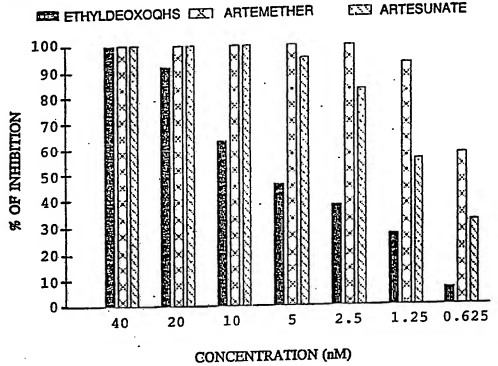
K1 strain from Kanchanaburi, Thailand (chloroquine resistant)

 $IC_{50} = 6.0 \text{ nM} = 2.0 \text{ ng/ml}$

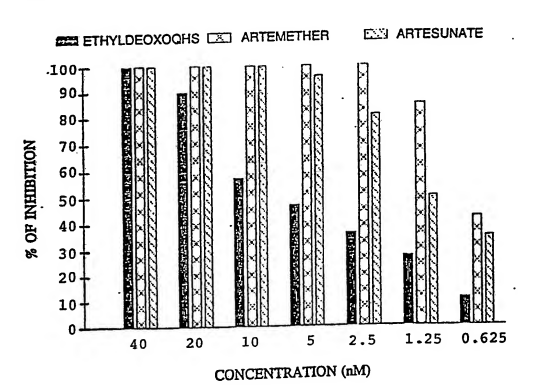
FC27 strain from Madang, Papua New Guinea (chloroquine sensitive)

 $IC_{50} = 4.3 \text{ nM} = 1.3 \text{ ng/ml}$

<u>In vitro</u> effects of ethyldeoxoqinghaosu, artemether and artesunate against the FC27 strain of <u>Plasmodium</u> <u>falciparum</u>



In vitro effects of ethyldeoxoqinghaosu, artemether and artesunate against the K1 strain of Plasmodium falciparum



Effect ginghaosu analogues on Tox lasma gondii in vitro*

	OHS Analogue	IC50 (μM)
	9 Deoxoqinghaosu	0.28
5	qinghaosu (natural compound)	0.9
	qinghaosu (prepared from qinghao acid)	0.85
	Dehydroqinghaosu sample 1	1.7
	Dehydroqinghaosu sample 2	1.7
	17 ß-epimer	2.2
10	19 ß-epimer	2.4
	19 α-epimer	4.4
	18 ß-epimer	4.8
	18 α-epimer	14
	Pyrimethamine (known compound)	1
·· 15	* [3H]-uracil incorporation assay	

Claims

1. Compounds of formula (I), pharmaceutically acceptable salts thereof or stereoisomeric forms thereof

$$H_3^{C} \xrightarrow{O-O} \begin{pmatrix} O-O \\ (X)_{N} \end{pmatrix}$$
(I)

5 wherein

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1 = 1, 2 or 3

n is an integer from 1-6

where X is independently selected from

H, =0,= CH_2 , aryl, COR, OR, COOR or

10 $X = -(CR_1R_2)_r - R_3$ where r is an integer from 1-10 and where r > 1, optionally at least 1 carbon atom can be replaced by 0, S or N;

 R_1 , R_2 and R_3 are independently selected from

aryl, each alkynyl, alkenyl, alkyl, H; one or by substituted optionally substituents selected from alkenyl, alkyl, alkynyl, aryl, halogen, OR, CF3, NO2, COOR, NRR', SR, COR, CONRR', SO3R, SO2NRR', SR, SOR, and SO2R, where R and R' are independently selected from H, optionally substituted alkyl, optionally alkenyl, optionally substituted substituted optionally substituted aryl alkynyl, optionally substituted arylalkyl wherein the optional substituents are as defined above; provided that $(C)_1-(X)_n$ is not $-CH_2-CH(CH_3)-.$

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- 2. A precess for preparing compounds of formula (I), pharmaceutically acceptable salts thereof or stereoisomeric forms thereof comprising:
 - (A) carrying out at least one addition reaction on a compound of formula (II) having a -C=0 functionality

$$H_{3}C$$

$$(C)_{1}$$

$$(V)_{r}$$

$$(V)_{r}$$

to provide another compound of formula (II) with an alcoholic functionality at the (C)₁-(V)_r group and then followed by oxygenation to give a compound of formula (I); where V is as defined for X; r and l are as defined above;

- directly oxygenating a compound of formula (II) (B) where there is a carboxylic functionality at the $(C)_{l}$ - $(V)_{r}$ reactive chain and also a facilitate subsequent functional group to compounds addition reaction, to provide formula (I) and then carrying out a suitable addition reaction to provide the required side chain; or
 - (C) for compounds of formula (I) where l=1; carrying out one or more of the following steps on a compound of general formula (II):
 - (i) oxidative degradation and
 - (ii) reduction

followed by oxygenation to give compounds of formula (I).

- 3. A process according to claim 2 wherein the oxygenation step in process A, B or C comprises
- 5 (i) initial irradiation under oxygen in the presence of a catalyst such as Rose Bengal and
 - (ii) further oxygenation by treatment with a transition metal catalyst.
- 10 4. A process according to claim 3 wherein the transition metal catalyst is one or more selected from Cu(OSO₂CF₃)₂, copper(II) carboxylic salts and iron(III) salts.
- 5. A process according to claim 4 wherein the Cu(II) carboxylic salt is Cu(II) propionate or copper(II) 2-ethylhexanoate and the iron(III) salt is Fe(phenanthroline)₃(PF₆)₃.
 - 6. A process according to claim 5 wherein process step A comprises
- 20 (i) treatment of the aldehyde 2 with A-MgBr, where A is alkyl, aryl or alkenyl to provide the respective derivatives; followed by oxygenation to provide a compound of formula (I);

with MeMgI to provide

followed by oxygenation to provide a compound of formula (I); or

- (iii) treatment of the aldehyde 2 with (EtO)₂CH-A-MgBr where A is alkyl;
 followed by hydrolysis and then oxidation to the HO(O=)C-A derivative;
 followed by oxygenation to provide a compound of formula (I).
- 10 7. A process according to claim 5 wherein in process B the starting compound of formula (II) is dehydroqinghao acid which is oxygenated to provide dehydroqinghaosu and the addition reaction is carried out by treating dehydroqinghaosu with thiol nucleophiles.
- 15 8. A process according to claim 7 wherein the thiol nucleophile is thiophenol, thioglycolic acid, methyl thioglycolate, mercaptopropionic acid, methyl mercaptopropionate, thioglycerol, L-cysteine or N-acetyl-L-cysteine.
- 20 9. A process according to claim 5 wherein process (C) comprises
 - (i) oxidative degradation of aldehyde 2
 - (ii) reduction to the alcohol
- (iii) followed by oxygenation to provide a compound of formula (I).

10. Compounds of general formula (II)

$$H_{3}C$$

$$(C)_{1}$$

$$(V)_{p}$$

wherein V is as defined for X; r and l are as hereinbefore defined; provided that the $(C)_1$ - $(V)_r$ group is not CH(CH₃)CH₂OH and CH(CH₃)C(=O)H.

- 11. A pharmaceutical composition comprising a compound of formula (I), a pharmaceutically acceptable salt thereof or stereoisomeric forms thereof in a pharmaceutically acceptable carrier and/or diluent.
- 10 12. A method of treatment or prophylaxis of parasitic or viral diseases in a mammal comprising administering to the mammal a compound of formula (I), a pharmaceutically acceptable salt thereof or a stereoisomeric form thereof.
- 13. Use of a compound of formula (I), a pharmaceutically acceptable salt thereof or a stereoisomeric form thereof in the manufacture of a medicament for the treatment or prophylaxis of parasitic or viral diseases.
 - 14. A compound of formula (I) substantially as herein described with reference to any one of examples 9-21 or 24.

In another aspect the present invention provides a process for preparing compounds of formula (I), pharmaceutically acceptable salts thereof or stereoisomeric forms thereof comprising:

5 (A) carrying out at least one addition reaction on a compound of formula (II) having a -C=0 functionality

to provide another compound of formula (II) with an alcoholic functionality at the (C)_I-(V)_r group and then followed by oxygenation to give a compound of formula (I); where V is as defined for X; r and l are as defined above;

- directly oxygenating a compound of formula (II) (B) 15 where there is a carboxylic functionality at the $(C)_{1}$ - $(V)_{r}$ reactive chain and also a further subsequent facilitate functional group to provide compounds of reaction, to addition formula (I) and then carrying out a suitable 20 addition reaction to provide the required side chain; or
 - (C) for compounds of formula (I) where l=1; carrying out one or more of the following steps on a compound of general formula (II):

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oxidative degradation and

(ii) reduction

followed by oxygenation to give compounds of formula (I).

Best Mode of Carrying out the Invention

basically, the process oxygenation step is The that described in PCT/AU90/00456 (WO 91/04970) and disclosure is incorporated herein by reference. oxygenation is preferably carried out as a "one-pot" reaction and involves oxygenation of a compound of formula (II) to provide a hydroperoxide derivative and without isolation further oxygenation in the presence of one or more oxygenating metal catalysts to give a of The oxygenation the compound of formula (I). hydroperoxy compound in the presence of one or more catalysts provides an oxygenation-cleavage-cyclization reaction to give the cyclic peroxyacetal compounds of formula (I).

oxygenation-cleavage-cyclization reaction The typically carried out by treating with one or more 20 transition metal catalysts such as Cu(OSO2CF3)2, Cu(II) propionate, copper(II) 2-ethylhexanoate, other copper(II) carboxylic salts, and various iron(III) salts such as Fe (phenanthroline) $_3$ (PF₆) $_3$. Other catalysts that may be used are cobalt(II) and cobalt(III) salts. 25 this reaction is carried out in a solvent such as a mixture of acetonitrile and dichloromethane treating with one of the above mentioned catalysts, or with a combination of the copper and iron catalysts. Other suitable solvents include hexane, ethyl acetate and 30 the like.

Preferred solvents for the Grignard addition

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PCT/AU92/00548

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reactions are diethyl ether, or THF but any other solvent such as benzene or other ether solvent would be suitable.

The reaction is typically initially carried out at the temperature of $.0^{\circ}-5^{\circ}\text{C}$ and continued at room temperature.

Other addition reactions are typically carried out in $CHCl_3$ and CH_2Cl_2 with an amine base. MeOH and aqueous MeOH may also be used and bases such as Na_2CO_3 , K_2CO_3 or $NaHCO_3$ can also be used. The reactions are preferably carried out at room temperature.

Oxidative degradation is preferably carried out using cupric acetate, 2,2'-bipyridyl and DABCO in DMF under atmospheric oxygen at between about 70°-75°C for about 12 hours.

The reduction is preferably carried out in methanol or other alcoholic solvent with NaBH₄ at about O^oC. Other reducing agents with appropriate solvents may also be used.

Preferably the starting compound for process A is the aldehyde 2 having the following structure

This is preferably treated with Grignard reagents or other organometallic reagents such as those derived from lithium, copper or zinc to provide compounds of the following formulae:

which are then oxygenated to provide the corresponding compounds of formula (I):

The aldehyde is preferably prepared from qinghao acid which has the following formula:

Qinghao acid occurs to the extent of 1-3% (dry weight) in Artemisia annua, which is much greater than the natural occurrence of qinghaosu and is easily extracted from the plant.

The starting material for the process B is preferably

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qinghao acid which is oxygenated to provide dehydroqinghaosu of the following formula

on which is carried out various addition reactions to provide compounds of formula (I).

The starting material for process C is also preferably the aldehyde having the structural formula 2. This aldehyde is typically prepared from qinghao acid by carrying out the following steps.

- methylation to provide the unsaturated methyl ester;
 - 2. reduction to provide the saturated methyl ester;
 - reduction to provide the alcohol;
 - oxidation to provide the aldehyde.

The skilled addressee would understand that process of the invention may result in one or more 15 stereogenic (chiral) centres being formed resulting in Thus it is to be understood that the stereoisomers. the its within scope includes invention present preparation of stereoisomers and also encompasses any isomers per se or mixture thereof. 20

Most of the starting compounds of formula (II) are also novel and form part of the present invention. Accordingly, in yet another aspect, the present invention provides compounds of general formula (II)

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qinghao acid which is oxygenated to provide dehydroqinghaosu of the following formula

on which is carried out various addition reactions to provide compounds of formula (I).

The starting material for process C is also preferably the aldehyde having the structural formula 2. This aldehyde is typically prepared from qinghao acid by carrying out the following steps.

- 1. methylation to provide the unsaturated methyl ester;
- 2. reduction to provide the saturated methyl ester;
- reduction to provide the alcohol;
- 4. oxidation to provide the aldehyde.

The skilled addressee would understand that the process of the invention may result in one or more stereogenic (chiral) centres being formed resulting in stereoisomers. Thus it is to be understood that the present invention includes within its scope the preparation of stereoisomers and also encompasses any isomers per se or mixture thereof.

Most of the starting compounds of formula (II) are also novel and form part of the present invention. Accordingly, in yet another aspect, the present invention provides compounds of general formula (II)

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